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This report supercedes and makes obsolete two earlier technical reports (Parts I and II) submitted to ARO entitled: "Studies on the Conversion of Pentaborane(9) to Decaborane (14)". We now have a simple method for producing in the laboratory good quality $B_{10}^{\dagger}H_{14}^{\dagger}$ from $B_{5}^{\dagger}H_{0}^{\dagger}$ in yields of 42-45%. Additionally we have been able to replace the costly reagents KH, BBr, and N(p-C,H₀), Br with the much cheaper reagents NaH, BCl₂, N(CH₂), Cl. We believe that this procedure is ready for scale up tests in $\frac{1}{2}$ to 1 pound quantities in a pilot plant reactor. Given below is a detailed description of our laboratory procedure for preparing

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Technical Report

STUDIES ON THE CONVERSION

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PENTABORANE(9) TO DECABORANE(14)

PART III

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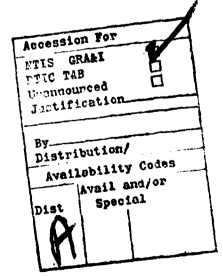
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Mark A. Toft



INTRODUCTION

This report supercedes and makes obsolete two earlier technical reports (Parts I and II) submitted to ARO entitled: "Studies on the Conversion of Pentaborane(9) to Decaborane(14)". We now have a simple method for producing in the laboratory good quality $B_{10}^{H}_{14}$ from $B_{5}^{H}_{9}$ in yields of 42-45%. Additionally we have been able to replace the costly reagents KH, BBr₃ and $[N(\underline{n}-C_{4}^{H}_{9})_{4}]$ Br with the much cheaper reagents NaH, BCl₃, $[N(CH_{3})_{4}]$ Cl. We believe that this procedure is ready for scale-up tests in $\frac{1}{2}$ to 1 pound quantities in a pilot plant reactor.

Given below is a detailed description of our laboratory procedure for preparing $B_{10}H_{14}$ in 1 gram quantities. It should be recognized that the laboratory method employs glass reactors and vacuum line techniques which are convenient for relatively small scale preparations. Some of the procedures employed in the laboratory method would be obviated in a scale-up procedure in which a metal reactor and metal piping would be employed. Thus, for example, while we condense B_5H_9 into a small glass reactor under vacuum at -196°, in a large scale procedure liquid B_5H_9 would be forced from its storage cylinder into the metal reactor by N_2 gas. In the stage where we maintain the mixture of BCl $_3$ and $[N(CH_3)_4][B_9H_{14}]$ at 11°, this is done to avoid excessive pressure in the glass reactor. With the metal reactor, room temperature would be permissible. Although vacuum sublimation is used to isolate $B_{10}H_{14}$ in the laboratory method, in the scaled-up procedure a stream of N_2 gas could be passed over the subliming $B_{10}H_{14}$ at 110° and a water-cooled collector would be sufficient to trap the $B_{10}H_{14}$.

A CONVENIENT LABORATORY PREPARATION

OF $B_{10}^{H}_{14}^{FROM}$ $B_{5}^{H}_{9}$

This procedure consists of three distinct steps or stages.

- 1. Conversion of B_5H_9 to $[N(CH_3)_4]B_9H_{14}$ by reacting a 2:1 molar ratio of B_5H_9 with NaH in THF in the presence of $[N(CH_3)_4]C1$ at room temperature.
- 2. After completion of step 1, THF is removed from the system and BCl_3 is added to the solid $[N(CH_3)_4][B_9H_{14}]$ and allowed to react at 11°.
- 3. After completion of step 2, $B_{10}H_{14}$ is sublimed at 110° from the solid reaction mixture.

Details of each of these steps follow.

1. 513 mg NaH (21.4 millimoles), 2.30 g [(CH $_3$) $_4$ N][C1] (22 millimoles), and a magnetic stirring bar ($1\frac{1}{2}$ " long) are placed in a 500 ml roundbottom flask in a dry box using N $_2$ atmosphere. The 500 ml flask is single neck with a 15 mm solv-seal joint glassblown in place of the normal neck opening. A reaction vessel to vacuum line stopcock adapter is attached to the 500 ml flask via this 15 mm solv-seal joint. The adapter is comprised of a 0 to 4 mm bore Kontes teflon stopcock to which is attached on one side a 15 mm solv-seal joint and on the other side a standard taper 14/35 outer joint.

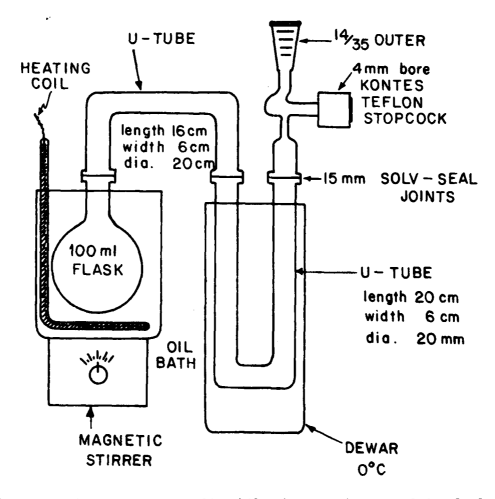
The reaction vessel (500 ml flask and stopcock adapter) is then removed from the dry box and attached to an inlet port of a standard vacuum line. The reaction vessel is evacuated of dry box N_2 and 20 ml of dry THF is condensed into the reaction vessel which is maintained at -78°C using a dry ice/isopropanol slush. The dry ice/isopropanol slush is then replaced with a liquid N_2 dewar and 4.10 ml B_5H_9 (42.8 mmoles), the volume measured at 0°C where the density of B_5H_9 is .66 g/m is then condensed into the reaction vessel. The reaction mixture is then warmed to room temperature (26-27°C) and is stirred vigorously for 12 hours. Immediate gas evolution is present and continues for several minutes.

The solution gradually changes from a clear slurry to light yellow. After 12 hours the reaction vessel is cooled to -196°C and the $\rm H_2$ produced is measured via a calibrated toepler system. 22 millimoles of $\rm H_2$ is obtained and this corresponds to initial deprotonation of one equivalent of $\rm B_5H_9$ with one equivalent of NaH. The reaction vessel is then warmed to 0°C while the solvent is removed under vacuum by pumping through a U-trap maintained at -78°C and a U-trap maintained at -196°C. Total solvent removal takes 3 hours, with the reaction vessel warmed to room temperature after 2 hours. The solid remaining is very light yellow and consists of [(CH₃)₄N][B₉H₁₄] and NaCl. ¹¹B NMR at 32.1 MHz shows the only boron containing species to be $\rm B_9H_{14}^{-1}$.

2. The $[(CH_3)_4N][B_9H_{14}]$ from step 1 is then scraped from the 500 ml reaction vessel and transferred to a similarly equipped 100 ml reaction vessel in the dry box. The stir bar is also transferred. Only a few, <10 mg are lost in the transfer. NOTE: The solid transferred also contains NaCl.

The 100 ml reaction vessel is removed from the dry box, placed on an inlet port of the vacuum line and evacuated of dry box N_2 . 22.0 mmoles of BCl₃ (freed from HCl) measured by gas volume at 26°C are condensed into the reaction vessel maintained at -196°C. The reaction vessel is then warmed to 11°C (H_2O and small quantity of ice) and stirred.

3. After 4 hours the reaction mixture is cooled to -196°C and 11 mmoles H_2 is removed using the aforementioned toepler system. The reaction vessel is then warmed to room temperature and traces of volatile materials are removed by pumping for a few minutes. The reaction vessel is then removed to the dry box where it is attached to a special "sublimator" for removal of $B_{10}H_{14}$. The "sublimator" system used is diagrammed below.



The U-tube is maintained at 0°C while the reaction vessel is slowly warmed to 110°C. The entire system is under dynamic vacuum. Decaborane collects slowly in the bridge and finally in the U-tube over a 2-hour period. Initially the $B_{10}^{\rm H}_{14}$ obtained is white but eventually is tinted light yellow by reaction by-products blown through the system due to dynamic vacuum. The amount of inpurity is however, quite small (a few milligrams). The initial mass of $B_{10}^{\rm H}_{14}$ obtained is 1.18 g or 45.2% based on $B_5^{\rm H}_9$. This is slightly reduced upon recrystallization which yields much purer $B_{10}^{\rm H}_{14}$ in a 1.10 g quantity or 42.1% yield based on $B_5^{\rm H}_9$ used. The $^{11}_{\rm B}$ NMR of the $B_{10}^{\rm H}_{14}$ shows it to be quite pure.